Pagetoid Reticulosis: Report of Two Cases and Review of the Literature

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Cutaneous T-cell lymphoma · Pagetoid reticulosis · Woringer-Kolopp disease · Mycosis fungoides · Skin tumors

Abstract
Pagetoid reticulosis is a rare variant of mycosis fungoides that presents with a large, usually single, erythematous, slowly growing scaly plaque containing an intraepidermal proliferation of neoplastic T lymphocytes. Histopathologically, this disease has distinctive attributes. In this report, we present two cases of pagetoid reticulosis, compare its microscopic features to those of ‘classical’ mycosis fungoides, and provide a brief review of the pertinent literature.

Pagetoid reticulosis (PR; also known as Woringer-Kolopp disease) is a rare, distinctive variant of mycosis fungoides (MF). It usually presents in adults as a single, erythematous, slowly growing scaly or verrucous plaque that is typically found on the extremities [1]. We herein report two cases of PR and compare its histological features to those of classical MF.

Case 1
A 61-year-old woman presented with a verrucoid pink-tan plaque on the right heel, measuring 3 cm in diameter, which was subjected to an excisional biopsy. It had been present for approximately 1 year. There were no cutaneous lesions elsewhere, and the remaining physical examination also showed no abnormalities. The results of a hemogram and a biochemical blood survey were normal.

Histological analysis showed a well-demarcated, discrete area of hyperkeratosis and parakeratosis with epidermal acanthosis. The surface epithelium was permeated by an infil-
A 50-year-old man developed a 1-cm pearly white-tan lesion on the chest, which had evolved over approximately 3 years. In the previous 3–4 months, it had increased in size and began to burn and itch.

Histological examination of an excisional biopsy specimen showed epidermal acanthosis with marked exocytosis, creating a ‘sieve-like’ appearance. The intraepithelial lymphoid cells were atypical cytologically, and they contained occasional mitotic figures. Those elements manifested significant irregularities in the nuclear membranes, with nuclear hyperchromasia and pleomorphism, and the cytoplasm was amphophilic or vacuolated. A mixed, superficial, perivascular dermal inflammatory infiltrate was also present (fig. 1). Immunohistochemical studies showed diffuse lesional reactivity for CD3, but labeling for CD7 was significantly diminished. Intraepidermal lymphoid cells were nearly all reactive for CD4, with only scattered staining for CD8 (fig. 2).

The patient is currently well, with no additional cutaneous lesions 16 months after the diagnosis.
Discussion

PR was first described by Frederick Woringer and Pierre Kolopp in 1939, in Strasbourg, France [2]. The first documented case was that of a 13-year-old boy who had a lesion on his forearm that measured <1 cm in diameter and had been present for 6 years; it then enlarged quickly. Original clinical diagnostic considerations included those of MF, melanocytic nevus, and Paget’s disease; an intraepidermal lymphoid infiltrate was observed microscopically [3]. In 1973, Braun-Falco and colleagues assigned the designation of ‘pagetoid reticulosis’ to this condition because of the intraepidermal pagetoid distribution of lymphoid cells, which were thought to possibly represent T cells [3]. Those authors believed that PR was a disease entity distinct from MF because of its localized nature and more marked lymphoid epidermotropism. Revuz and colleagues later concluded that the Woringer-Kolopp disease and PR were one and the same condition [3].

Currently, it is still difficult to distinguish PR from other variants of MF by microscopy alone. Indeed, the differential diagnosis of PR includes MF of the ‘palmaris et plantaris’ type. Primary epidermotropic CD8+ cytotoxic T-cell lymphoma represents another consideration [4]. PR almost exclusively comprises dense intraepidermal lymphoid infiltrates with variable acanthosis [5]; few atypical cells are seen in the dermis [1], but the corium commonly contains mature lymphocytes and histiocytes. Immunologically, the lesional cells in PR are typically CD4+ T cells, although examples with CD8+ and CD4−/CD8− phenotypes have also been described [5]. The cellular proliferation rate, as measured immunohistologically with the Ki-67/MIB1 antibody, is usually >30%, and CD30 reactivity may be present as well [5].

In comparison, classical MF typically manifests both epidermal and dermal components; the latter comprise a polymorphous cell population consisting of neoplastic and mature lymphocytes, histiocytes, plasma cells, and eosinophils [5]. Clinically, typical MF lesions are
most often thin plaques on the distal extremities and trunk [5]. Immunologically, the tumor cells are CD4+ with a low proliferation index (<10%), and they are usually CD30− [4]. Despite those differences, the World Health Organization has recently classified PR as a unilesional variant of MF [6].

PR is generally believed to be an indolent disease. However, it seems that this is not always the case. In a study by Haghighi et al. [5], some patients with PR had clinically progressive lesions that required aggressive therapy. Another publication suggested that it was sometimes difficult to distinguish PR from MF of the ‘palmaris et plantaris’ type, and that the latter cases could require active treatment [7].

It is important to include other aggressive T-cell lymphomas in the differential diagnosis of PR. For example, in one reported case, a patient with high-grade peripheral T-cell lymphoma was erroneously categorized initially as PR. The tumor subsequently exhibited visceral involvement and proved fatal in <2 years [8]. Another similar example was that of adult T-cell lymphoma/leukemia that was first thought to represent PR; it likewise pursued an aggressive clinical course [9].

Immunohistochemical analysis of the lesions and thorough clinical correlation are necessary to avoid such problems. Recent investigations of T-cell receptor gene rearrangements in PR have shown that either TCRγδ or TCRαβ may be aberrant [10]. The proliferative index of the lesional cells is only debatably related to prognosis [11].

Recently documented cases of PR [12–17] have included patients of all ages, including children. Localized perioral lesions have also been reported [18] as well as a generalized form of the disorder known as Ketron-Goodman disease [19]. On dermoscopy, PR has been said to show features that are similar to those of Bowen’s disease. They include glomerular and dotted vessels, whitish scaly areas, and reticular depigmentation [20].

Several treatments have been employed in PR cases, such as topical steroids, topical nitrogen mustard, light therapy, interferon alpha-2b [21], localized [22] and modulated [23] radiation therapy, imiquimod [24], altretinoin [25], and fractional laser-assisted photodynamic therapy [26, 27]. In the reported experience of Lee et al. [22], localized radiation therapy seemed to produce the best results. Photodynamic therapy may be an alternative to irradiation in young patients [27].

In summary, we have presented two cases of PR. Their features and a review of the literature support the conclusion that PR is a unilesional variant of MF with distinctive clinico-pathologic features.

Statement of Ethics

The approval of the authors’ institutional review board was obtained for this study as well as the patients’ consent.

Disclosure Statement

The authors have no fiduciary interests or conflicts of interest to disclose.
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